(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 30.08.2000 Bulletin 2000/35

(21) Application number: 99103814.2

(22) Date of filing: 26.02.1999

AL LT LV MK RO SI

(51) Int CI7: CO7D 213/24, CO7D 213/38, CO7D 213/53, CO7D 213/89, CO7D 401/12, A61K 31/44, A61K 31/495, A61K 31/55, C12Q 1/68

(84) Designated Contracting States: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE Designated Extension States:

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(54) Inhibitors of cellular nicotinamide mononucleotide formation and their use in cancer therapy

(57) New biologically active substances are described which inhibit the cellular formation of niacinamide mononucleotide, an essential intermediate of the NAD(P) biosynthesis in the cell. These substances can represent the active incredient of a pharmaceutical.

composition for the treatment of cancers, leukaemias or for immunosuppression. Furthermore, screening methods are described as a tool for delecting the above active substances, and for examination of a given cell type for its dependency on niacinamide as a precursor for NAD synthesis.

Description

Fleid of the invention

[0001] The present invention relates to new biologically active substances which inhibit the cellular formation of niacinamide mononteolotide, which is one of the assential intermediates in the NAD(P) biosynthesis in the cell. The invention further concerns pharmaeutical compositions containing these substances and their use, especially in the treatment of cancers, leukaemias or for immunosuppression. The invention also provides screening methods as a tool for detecting the above active substances, and for examination of cell types with respect to their NAD(P) synthesis pathway.

Technical Background of the Invention

[0002] NAD is synthesized in mammalian cells by three different pathways starting either from tryptophan via quinolinic acid. from niacin (also referred to as nicotinic acid) or from niacinamide (also referred to as nicotinamide), as shown in Floure 1.

[0003] The addition of a phosphoribosyl moiety results in the formation of the corresponding mononucleotides, niacin mononucleotide (NAM), Quinclinic acid is utilised in a reaction with phosphoribosyl pyrophosphate (PRPP) to form niacin mononucleotide (NAM). The enzyme catalysing this reaction, quinclinic acid phosphoribosyl transferase (©), is found in liver, kidney and brain.

[0004] Niacin reacts with PRPP to form niacin mononucleotide (dNAM). The enzyme catalysing this reaction is niacin phosphoribosyl transferase (②) and is widely distributed in various tissues. Both pathways starting either from tryptophan or from niacin as NAD precursors merge at the stage of the niacin mononucleotide formation.

[0005] Niacinamide reacts with PRPP to give niacinamide mononucleoide (NAM). The enzyme that catalyses this reaction is niacinamide phosphrorbooyl transferase (a). This enzyme is specific for niacinamide and is entirely distinct from niacin phosphorbooyl transferase (b). It is also widely distributed in various tissues.

[0006] The subsequent addition of adenosine monophosphate to the mononucleotides results in the formation of the corresponding dinucleotides. Nacin mononucleotide and niacinamide mononucleotide react with ATP to yield niacin adenine dinucleotide (nNAD) and niacinamide adenine dinucleotide (NAD) and niacinamide adenine dinucleotide (NAD) prophosphorylase (0).

But the different pathways, are calalysed by the same enzyme, NAD pyrophosphorylase (0).

[0007] A further amidiation step is needed to convert fination adentine dinucleotide (dNAD) to niacinamide adenine dinucleotide (NAD). The enzyme which catalyses this reaction is NAD synthetase (**9**). NAD is the immediate procursor of niacinamide adenine dinucleotide phosphate (NADP). The neaction is catalysed by AND kinase (6). For details see, for example, Covy, J.G. Purine and pyrimidine nucleotide metabotism. In: Taxbook of Biochemistry and Clinical Cornelations, 3rd edition, ed. Devitin, T, Wildy Brisbane 1992, pp529-574.

[0008] Normal cells can hypically utilize both precursors niscin and niscinamids for NAD(P) synthesis, and in many cases additionally hypiophan or is metabolites, which has been demonstrated for various normal tissues: Accordingly, Murine glial cells (cortex and hippocampus = brain) use; niscin niscinamide, and quindinic acid (Grand et al. (1995), J. Neurochem, 70: 1759-1763), Human hypincoyles use niscin and niscinamide (Carson et al. (1997), J. Immunol, 138: 1904-1907). Barger at al. (1992), Ep. Cell Res, 177: 79-8B), Ratt ver cells use niscin, niscinamide and trylprophan (Yamada et al. (1993), Internat. J. VII. Nutr. Res, 53: 184-191; Shin et al. (1995), Internat. J. VII. Nutr. Res, 56: 142-145; Ostrich (1971), Mehrods Enzymin 188: 144-149, Human eyphrocytes use niscin and niscinamide (Rockingiani et al. (1991), Purine and privrieting and privriet

5 (0009] NAD(P) is involved in a variety of biochemical reactions which are vital to the cell and have therefore been thoroughly investigated. This key function of NAD(P) has evoked also some investigations in the past on the role of this compound for the development and growth of tumors, and as to what the NAD(P) metabolism could also be utilized to combat tumors. Indeed, substances aiming at the treatment of tumor diseases then been described which involved concentrating the contraction of the national contraction of the properties of the properties of diseases of NAD(P) levels in the cell. However, these substances primarily of act by initiating the cellular synthesis of disuclectife derivatives which structurally divide from natural NAD. The blockholms of the properties of the properties of the properties of the resulting cell-damage are, therefore, manifold as outlined in the Table 1.

Table 1

Compounds	Mode of action	Ref
6-aminonicotinamide	Primary mechanism of action:	
	Synthesis of 6-amino-NAD(P), a competitive inhibitor of NAD(P)-	1, 2, 3
	requiring enzymes, especially of 6-phosphogluconate	1
	dehydrogenase, an enzyme of the pentose-phosphate-pathway	
	which provides the precursor of the nucleotide biosynthesis	1
	ribose-5-phosphate.	
	Resulting blochemical effects in the cell:	
	Inhibition of purine nucleotide de novo synthesis from	1
	[14C]glycine.	1
	Decrease of intracellular purine (ATP, GTP) and pyrimidine	
	(UTP, CTP) nucleotide pools resulting in the inhibition of DNA	
	and RNA synthesis.	
	Inhibition of PARP (an enzyme involved in the DNA repair).	
	Reduction of the ATP to ADP ratio.	
	Depression of intracellular NAD concentration,	
tlazofurin, selenazofurin	Primary mechanism of action:	
	Synthesis of the NAD analogs TAD, SAD which are potent	2, 4, 5
	inhibitors of inosine monophosphate dehydrogenase, an enzyme	1
	involved in the synthesis of purine nucleotides.	
	Resulting biochemical effects in the cell:	
	Depletion of GMP and accumulation of IMP resulting in an	1
	inhibition of DNA and RNA synthesis.	1
	Stimulation of NAD synthesis after short exposure (<24 h).	1
	Inhibition of NAD synthesis after prolonged exposure	1
	(>24 h), probably due to negative feedback inhibition of NAD	
	synthesis by TAD/SAD which accumulate in the cell.	
azaserine,	Primary mechanism of action:	
6-diazo-5-oxo-L-	Analogs of glutamine which block the enzymatic transfer of the amido	6, 7
norleucine	group of glutamine.	
	Resulting blochemical effects in the cell:	
	Inhibition of IMP synthesis resulting in an inhibition of DNA and	
	RNA synthesis.	
	Inhibition of NAD synthesis from the precursor niacin at the	
	following step: dNAD -> NAD	l
	Tollowing stop. Stand - 1400	
	Mutagen, cancerogen.	
DNA-Interacting compounds	Primary mechanism of action:	
compounds	Induction of DNA strand breaks.	4, 8, 9,
	INDUSTRIA STATE STATE OF STATE	4, 0, 8
		l

Table 1 (continued)

Compounds	Mode of action	Ref.
(e.g. N-methyl-N'-nitro-	N-methyl-N'-nitro- Resulting blochemical effects in the cell:	
N-nitroso-guanidine)	Multiple consequences of DNA damage.	
	Activation of the DNA repair enzyme PARP resulting in a decline	
	of the intracellular NAD content, since the substrate of PARP is	
	NAD.	
	Mutagen, cancerogen.	
Abbreviations: PARP	poly(ADP-ribose) polymerase; NAD, niacinamide adenine dinucleotide:	
NADP, niacinamide adenin	e dinucleotide phosphate; dNAD, niacin adenine dinucleotide; ATP,	
	DP, adenosine diphosphate; GTP, guanosine triphosphate; GMP,	
	; UTP, uridine triphosphate; CTP, cytosine trisphosphate; DNA,	
desoxyribonucleic acid; RN	A, ribonucleic acid; TAD, tiazofunn adenine dinucleotide; SAD.	

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[0010] It is therefore not possible to make any predictions from these data on the biological effects of a primary and specific inhibition of the NAD biosynthesis in various cell types. In particular, it remains completely speculative whether this mechanism may be advantageous over the above utilisation of diructeotide derivatives with regard to tumor selectivity of the cell damaging effect, the most important feature of a potential drug for tumor therapy.

selenazofurin adenine dinucleotide; IMP, inosine monophosphate.

[0011] JP-459555, published in 1970, describes the extraction of a structurally unknown constituent from potatoes. baker's yeast and bovine blood which inhibits respiration of tumor cells and NAD synthesis of enythrocytes. The inventors propose the use of this constituent for tumor therapy. However, the data presented in JP-459555 are far from making it clear or even probable that inhibition of NAD synthesis is useful for the therapy of cancer. The inventors rather come to the conclusion that the biological activity of the substance is multifaceted and not limited merely to the phenomenon of inhibition of NAD biosynthesis. In a study published later by the same research group (A. Kizu: Kvoto Furitsu Ika Daigaku Zasshi 80, pp. 14-24, 1971) it was shown that the extracted compounds (derivatives of glucose) inhibit respiration and glycolysis in tumor cells already within a few minutes in addition to inhibition of NAD synthesis. In fact, tumor cells treated with the extract for only 20 min suffered from such heavy damage that they did not grow in the abdominal cavity of mice in contrast to untreated control cells. In contrast to this finding, the present inventors have observed that compounds which promptly and selectively inhibit NAD synthesis in the cell exert a deleterious effect on tumor cells only after an exposure for 3-4 days, whereas an exposure for 20 min is completely ineffective irrespective of the concentration employed. Thus, it is very unlikely that it is NAD biosynthesis inhibition by which the extract disclosed in JP-459555 damages tumor cells. It is rather assumed that other mechanisms are primarily responsible for the cell death, while the reduction of the NAD levels is a secondary effect due to the general damage to the cell. The prompt deleterious effect on tumor cells as produced by this extract is, therefore, obviously due to a inhibition of cell

[0012] Also, the tumor preference of the cell killing effect of the extract, as described in JP-459555, can easily be explained by a characteristic feature of the respiration inhibiting effect of the extract, as this effect is marked in tumor cells but absent in liver cells. (A. Kizu, Figure 2). Thus, clearly JP-459555 did not disclose any means to affect tumor cells by NAD synthesis inhibition.

[0013] It was also known that DNA damaging cytotoxic compounds initiate a decrease of the cellular concentration of NAD. Some authors assumed that lowering of cellular MAD levels, with a resulting shortage of ATP within the cell, may play a role in the mechanism of cell death produced by these compounds (Daniel S. Martin and Gary K. Schwartz, Oncology Research, Vol. 9, pp. 1-5, 1997). The effect of these compounds on the NAD concentration within the cell ersults, however, indirectly from an enhanced NAD consumption by enzymes involved in DNA repair (see Table 1). [0014] The primary effect of these compounds, namely damage to the DNA, has many consequences in addition to lowering cellular NAD levels. As known, the DNA is in control of the synthesis of many cellular constituents, like proteins and enzymes, which are of vital importance to the cell. Thus, the consequences of DNA damage are also manifold, lowering of the cellular NAD concentration being only one of them. The efficacy profile of a specific inhibition of the NAD biosynthesis can, therefore, not be concluded from observations made with these compounds.

[0015] Just as title information on what can be expected from a specific inhibition of the biosynthesis of NAD gives the symptomatology of niscinarnide and niscin deficiency. These vitamins of the B group are procurous of the NAD biosynthesis as outlined above. Long term deficiency of these precurous results in a disease known as pellagra. Main symptoms are alterations of the skin and dementia. This syndrome shows no similarity to the chronic intoxication with any of the compounds discussed above.

[0016] Thus, in summary, the state of the art does not allow to draw conclusions as to what can be expected from a

primary and specific inhibition of the cellular synthesis of NAD because the compounds known to lower the cellular. NAD concentration exact other primary effects which may affect cell survival by thereselves. There exists no other reliable means to solve this question than the use of a specific inhibitor of NAD synthesis. But no such compound was available in the past:

5 (0017) Morton (R. K. Morton: Nature 181, pp. 540-543, 1958) proposes for human cancer therapy to aim at compounds which inhibit the NAD pyrophosphorylase (Enzyme 0) in Figure 1) as the activity of this enzyme was assumed to be the limiting factor of NAD synthesis. Note that the biosynthesis pathway from both niacin and nilacinamida, and also from tryptophan would be blocked by an inhibition of the NAD pyrophosphonylase since it acts on a late step of the biosynthesis pathway where the initiality separated pathways starting from the different precursors tryptophan capical or nicerial conduction of the NAD pyrophosphonylase since it acts on a late step of the biosynthesis pathway where the initiality separated pathways starting from the different precursors tryptophan conduction in low Thus no evidence for the correctness of this assumption is available for this enzyme has been found until mor. Thus, no evidence for the correctness of this assumption is available.

Detailed Description of the Invention

- 5 [0018] The present invention is based on the surprising finding that specific cell types can essentially utilize only niacinamide as a precursor for the cellular NAD(P) biosynthesis. Niacin or typtophan which constitute alternative precursors in all other cell types investigated so far cannot, or at least not to a significant extent, be utilized. Accordingly, the present invention provides for biologically active substances which inhibit the cellular formation of niacinamide mononucleotids. Substances having this activity, can easily be identified by the screening assay described below. Preferably, the present substances exhibit an inhibitory activity on cellular NAD biosynthesis from the precursor niacinated at concentrations of ≤ 10 µM of 50 %, more preferably 80 % and most preferably 90 % in such an assay. [0019] In this connection it is noted that inlacinamide, which has been taken up, for example, with the dief, has first to be channeled into the cell across the cell membran before it can be converted by niacinamide phosphoriboely transferase into inclaimantide connoclated into the reliablement of the present invention covers therefore not only hibitors of the insichamide
- brane.

 (Mith the compounds of the present invention it is possible for the first time to damage exclusively those cells which mainly use niacinamide as a procursor for NAD biosynthesis saving those cells which are additionally able to synthesize NAD form niacin or trylopchan (Figure 1). It tumed out that by using hese compounds many malignant cells are affected while non-malignant cells can be saved. The same applies to certain lymphocytes which play a role in immune reactions. This behaviour has not been observed before and is also completely surprising. Neither there was any indication in the prior at nor could it be expected on the basis of the known data that normal somatic cell which can typically use all three kinds of precursors loose their ability to accommodate tryptophan and niacin and become dependent only on nicknamide when turning malignant.

phosphoribosyl transferase, but also substances that hinder or block the transport of niacinamide across the cell mem-

- 35 [0021] Thus, biologically active compounds which selectively block the niacinamide branch of the NAD biosnihasis, i.e. inhibit the formation of niacinamide mononucleotide on the cellular level, would offer a new approach for selective tumor therapy. Melignant colls dependent on nicinamide as a main or sole precursor would suffer from such damage and finally be destroyed due to the inhibition of niacinamide mononucleotide formation and the subsequent NAD(P) operation. On the other hand, normal somatic cells can compensate for the inhibited niacinamide branch by still utilizing niacin and/or tryotophan as or excursors thereby ovovition sufficient NAD levels to cusarate survivel of the cells.
- [0022] The compounds of this invention are the first which primarily and specifically inhibit the biosynthesis of NAD from niacinamids. Therefore, these compounds can be used as a tool for investigation on the effect of this primary event on the survival of tumor cells and other cells of the body.
- [0023] Moreover, it was completely surprising that the new specific inhibitors of NAD synthesis via ni acinamide which deplete NAD in turn or cells within hours did not quickly fell the cells as shown with the known NAD synthesis* inhibitors (see Table 1) but ratine produced a characteristic "delayed cell death" phenomenon in these cells: continued growth for up to 3 days was observed in presence of the new compounds before practically all cells underwent apopticit cell death. It was additionally surprising that many non-malignant cells are very resistant to the apopticies-indusing effect of the new specific inhibitors of NAD biosynthesis. For restance a 10000-10d higher concentration is necessary to kill burners bore marrow cells compared to most tested furnan cancer cell lines. Thus, the "delayed cell death" charges.
 - teristic can be used in an assay to screen for substances according to the present invention.

 [0024] The ability of the compounds of the invention to inhibit the ADD biosynthesis from nacinamide can be shown
 with an easily reproducible test system which measures the incorporation of radioactive niacinamide into NAD and

 NADP. This assay provides a further sreening system to examine any chemical substance for its ability to selectively
 inhibit the cellular niacinamide mononuclocitide formation. The assay allows to screen for and select the inhibitory
 compounds of the present invention without being bound to a particular structural characterization. Accordingly, there
 is no limitation on the chemical structure of these compounds as long as they exhibit is side specific inhibitory activity.

and any known preparation methods can be used.

[0025] The fact that the death of tumor cells initiated by these compounds is indeed due solely to the inhibition of the NAD Diseynthesis from niscinsmide and not due to any other effect could be unequivecally verified. Addition of excess niscinamide to the astracellular medium in which the cells are grown in vitro completely reverses the apoptosis inducion effect of the new compounds.

[0026]. The effects of the compounds according to the invention on cell growth under high-density conditions have been investigated in order to closely simulate the *in vivo* situation of solid fumors. For this purpose, the inventors seeded high cell numbers and carried out high density cell culture experiments with the compounds shown in Table 2 below. Cell growth was monitored at various times up to 10 days as described in the experimental part below. Human hepatocarcinoma cells (HaoC2) were used. for example.

[0027] The time curve of the action of the compounds is characterised by the induction of "delayed cell death" which is clearly distinguished from a rapid decline of cell numbers occurring after the application of toxic compounds. The "delayed cell death" phenomenon is described using for example the results obtained with K22-339. Figure 2 demonstrates the characteristic time curve of growth inhibition by substance K22-339. During incubation with K22-339 the number of Hop62 cells increased up to three days, after which the culture was no longer able to grow and cell numbers declined from day 7 to 10. Cell cleath occurred on day 4, and the cells gradually detached until day 10, in contrast, toxic compounds are lies sactive in high density cultures and effective concentrations funded a rapid decrease in cell numbers observed as soon as one to three days of incubation. Using K22-339, however, a concentration of 0.3 µM was sufficient to bring about the tuil-blown effect. A still ten innes higher concentration of an either brown excelled the time until the cell number gradually decreased. This characteristic action was referred to as delayed cell death.

Table 2

K-No.	Structure	DCD [µM]	
K22.132	of chimino	0.1	
	4-benzhydryl-piperazine-1-carboxylic acid-[6-(3-pyridine- 3-yl-methylureido)-hexyl]-amide		

K-No.	Structure	DCD [µM]
K22.234	1-[5-(1-benzhydryl-piperidine-4-yl)-pentyl]-3-pyridine-3-yl-thiourea	0.3
K22.265	gonito	3
	6-(4-benzhydryl-piperazine-1-yl)-hexanoic acid-(2- pyridine-3-yl-ethyl)-amide	
K22.299		0.3
	1-(6,6-diphenyl-5-hexenyl)-3-(pyridine-3-yl-methylene- amino)-thiourea	
K22.339		0.1
	1-[4-(1-benzhydryl-piperidine-4-yl)-butyl]-3-(2-pyridine-3-yl-ethyl)-urea	
K22.350	00	1
. 1	N-{2-[5-(4-benzhydryl-piperazine-1-yl-methyl)-1-methyl- 1H-pyrrole-2-yl]-ethyl)-3-pyridine-3-yl-acrylamide	

K-No.	Structure	DCD [µM]
K22.387	Sjon-j.jo	0.01
	1-(4-[1-(10,11-dihydro-dibenzene[b,f]azepine-5- carbonyl)-pipendine-4-yl]-butyl]-3-pyridine-3-yl-urea	
K22.408	3140	1
	2-amino-3-[4-hydroxy-3-(2-[4-[4-(3-pyridine-3-yl-acryloyl-amino)-butyl]-piperidine-1-carbonyl]-phenylazo)-phenyl]-propanoic acid trihydrate	
K22.130		1
	1-[4-(1-benzhydryl-piperidine-4-yl)-butyl]-3-pyridine-3-yl- urea	
K22.158		1
	1-(3,3-diphenylpropyl)-3-[6-(3-pyridine-3-yl-methylureido)-hexyl]-urea	
K22.316	orange o	1
	N-(4-(1-[4-(1-benzhydryl-piperidine-4-yl)-butyl]-piperidine- 4-yl]-butyl)-3-pyridine-3-yl-propanoic acid amide	

K-No.	Structure	DCD [µM]
K22.365	Similar .	3
	1-{4-[1-(naphthalin-2-sulfonyl)-piperidine-4-yl)-butyl}-3- pyridine-3-yl-urea	

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[0028] The DCD-value was defined as the minimum concentration of the respective compound, which - despite initial growth of the culture - induced cell death blow the number of initially seeded cells. All compounds were active on high density cultures of HepG2 cells at concentrations of 3 μ M or lower. It is therefore preferred that the compounds of the present invention are active in the "delayer cell delath" test at a concentration of 3 μ M or lower, particularly preferred at a concentration of 1 μ M or lower. The compounds listed in Table 2 were prepared according to standard methods known in the art.

© [0029] The time course of the effect of the compounds suggested that there was no acute unspecific tocicity on the tumor cells. This time course is in sharp contrasts to the results described in JP445855. The authors wore that a 20 min incubation period of tumor cell with the disclosed extract was sufficient to induce cell death. On the other hand, the present compounds seem the imposes some physiological limitation on the cells which should leave enough time for them to sense the restriction and commit suicide before uncontrolled necrosic could take piece. "Delayed cell death" is induced by the compounds occurred in the form of apoptosis which is a favourable way of removing no longer viable cells, because it avoids the unregulated release of cell contents to the surrounding issue. The "delayed cell death" induced by the compounds could be antaconised by the addition of hischamida.

[0030] The effect of the compounds on the synthesis of NAD(P) starting from niacinamide was investigated using for example the tumor cell line HepG2 according to the technique described in the experimental part below. As shown in Table 3, the compounds completely inhibited the de nove synthesis of NAD(P) from its executes or niacinamide.

Toble S

latie 3						
K-No.	NAD(P) pmol/10 ⁶ cells	NAD(P) pmol/mg protein	% of Control			
Control	302.1	186.4	100			
K22.132	4.1	2.5	1.4			
K22.234	3.4	2.1	1.1			
K22.265	7.3	4.5	2.4			
K22.299	1.6	1.0	0.5			
K22.339	2.5	1.6	0.8			
K22.350	2.7	1.7	0.9			
K22.387	2.6	1.6	0.9			
K22.408	4.9	3.0	1.6			

[0031] The inhibition of the NAD(P) synthesis from niacinamide by these compounds was investigated in HepG2 cells as described in Material and Methods. The compounds were used at a concentration of 10⁵ M. As a control, which treated or untreated realls were used.

[0032] In these experiments, a pre-incubation period of 17 his with the compounds was used, but it turned out that he pre-incubations period outlobe is bentdered, for example to 2Ps, or skepped completely without affecting the inhibition profile. Further analysis of the radiolabelled niscinamide metabolities revealed that the compounds exclusively blocked he formation of the niscinamide monouncledoids. Figure 3 shows representative chromotograms of [145]-inscinamide metabolities extracted from vehicle - and K22 234-treated HepQ2 cells. Similar results werde obtained with the other compounds.

[0033] By comparing the two chromatograms in Figure 3, it is clearly evident that the compounds inhibited the synthesis of NAD(P) from its precursor neisonamics. The radiolabelled naich seen in the extract of the verbicle- and the compound-treated cells resulted from the enzymatic deamistation of [14C]placinamide. The peaks of NAD and niscin on the chromatograms are close together, but the inventors verified the identification by the second thin-layer chromatography system, as described in Materials and Methods. Since an accumulation of the intermediate niscinamide mononucleotide was not detected in the compound-treated cells, the compounds inhibit the enzyme niscinamide prophosphate phosphoribody transferase (EC 2.4.2.12); a second rame of this enzyme is niscinamide mononucleotide prophosphate phosphoribody transferase (EC 2.4.2.12); a second rame of this enzyme is niscinamide mononucleotide prophosphate phosphoribody transferase (EC 2.4.2.12); a second rame of this enzyme is niscinamide mononucleotide prophosphate (PC)placinamide as precursor related to the cell. Performing the same kind of experiments using (PC)placinamide as precursor raisent in cells which are able to use the niscin pathway. The observation that the nicin pathway of NAD(P) synthesis is not blocked by the compounds makes it very unlikely that pathway starting from tryptophan is suppressed by the compounds.

[0034] Before the advent of the present invention, no compounds have been described which exclusively inhibit the NAD(P) synthesis at this step. Accordingly, this mode of action is entirely different from that proposed by Morton (Nature 1815:65-643, 1959) for tumor therapy. He suggested to inhibit the extryme NAD pryproposphorylase, but he inhibition of this enzyme would block the synthesis of NAD(P) from all three precursors (niachamide, niacin and tryptophan, compare Pigure 1). It was alsolutely surprising that the inhibition of NAD synthesis at the step of the formation of niachamide mononculcedict in the niachamide behave was sufficient to kill most tumor cells.

[0035] The present compounds are the first tools to investigate the importance of the niacinamide pathway in tumor calls. The present inventors found that most tumour calls, for example, HopQ2 (liver carcinoma), U.P.M. (6) (globals-coma, astrocytoma), U.373 MS (globals-coma, astrocytoma), U.373 MS (globals-coma, astrocytoma), U.373 MS (globals-coma, astrocytoma), MCP.T.M (freest class call carcinoma), KG-la (myslogenic laukaemila), HL-Q0 (promejector) tealutemila, PSQ (lung carcinoma), MCP.T.M (freest carcinoma) (25) grosstate carcinoma), Can utilize only on the niacinamide pathway of NAD(P) synthesis. From this finding it is concluded that these compounds can be used for the tharapy of the corresponding cancers (e.g. breast, prostate, lung, colon, carvix, ovan, skin, CNS, bladder, pancreas and leukemia and lymphoma). The above cell lines are known and commercially available.

[0036] Furthermore, blocking only the niacinamide pathway protect those cells from the effect of the compounds which can also use niacin or tryptophan as precursor. These cells are typical healthy somatic cells, e.g. liver cells, Kupffer cells, lung epithelial cells, renal apithelial cells, lymphocytes, colon epithelial cells or dermal fibroblasts. In regard to side effects, this essentially sole inhibition of the niacinamide pathway is of enormous advantage for the treatment of canner.

[0037] For the detection of specific inhibitors of nacinamide mononucleotide formation the following screening assay is especially useful within comprises the following steps: incubating cultured calls selected from HegG2 calls, U87 5 MG calls, MCF-7 Mt calls, Cali-1 cells, IL-60 calls, PG2 calls, U873 MG calls, A549 calls and KG-1a calls in the presence of [1⁴C)-inacinamide and a substance to be tested for its activity to inhibit the callular comprused and measuring the amount of formed labelled niacinamide mononucleotide, NAD and NADP. More concretely, cultured calls are seeded in a defined density in culture defises, followed by incubation in the presence of a test compound and by the addition of [1⁴C)-inacinamide for about 0,1 to 6 hrs. The calls are then lysed with perchioric acid and the resulting extract is neutralised. Finally, the [1⁴C)-belied compounds are separated by thin-layer chromatography on cellulose matrices (followed by UV detection and autoradiography. Non-radioactive nilacinamide derivatives are used as standards. This assay allows employ and effective screening and selection of the compounds according to the invention.

[0038] The present invention also provides a method for assaying the dependency of a given cell type on niacinamide sa as a precursor for NAD synthesis. This method allows to delarmine which (malignant) call types are perticularly sensitive to the substainces according to the invention and can be helpful to develop a suitable regimen for combatting various tumors. Accordingly, such an assay would comprise inclusting cells to be I estad in the presence of a substance according to the invention in a medium containing only niacinamide as a NAD synthesis procursor, and performing a cytlotxicity assay after the incubation period. Such a cytotoxicity assay could, for example, be the "high desity cell of lest "described in the experimental part."

Example

Material and Methods

5 Reagents:

[0039]

Trypsin/EDTA: 0.05 % (w/v) trypsin (Difco, Detroit, USA) + 0.016 % (w/v) EDTA (Sigma, Deisenhofen,

Germany).

[14C]Niacinamide: ARC794, American Radiolabeled Chemicals Inc., St. Louis, MO, USA, 0.25 mCi/ml;

specific activity 50 mCi/mmol.

Lysis buffer: 0.5 M perchloric acid (Merck, Darmstadt, Germany).

Neutralisation reagent: 0.5 mM potassium chloride, 2.0 M potassium hydroxide, dissolved in purified water.

The chemicals were obtained from Merck, Darmstadt, Germany.

TLC foils: Cellulose F, Art. 1.05565, Merck, Darmstadt, Germany Poly(ethyleneimine) Cellulose F, Art. 1.05579, Merck Darmstadt, Germany.

TLC solvents: 0.05 M lithium chloride or 3 parts 1 M ammonium acetate pH 5.0 + 7 parts ethanol

(Merck, Darmstadt, Germany).

20 Standards: 10 to 20 mo/ml solutions of the following niacinamide derivatives were prepared: niacin.

10 to 20 mg/ml solutions of the following niacinamide derivatives were prepared: niacin, niacinamide, niacin mononucleotide (dNAM), niacinamide mononucleotide (NAM), niacinamide mononucleotide (NAM), niacinamide adenine dinucleotide (NAD), niacinamide adenine dinucleotide (NAD), niacinamide adenine dinucleotide (NAD), niacinamide adenine dinucleotide ohosohate (NADP). All standards were purchased from Sionamide adenine dinucleotide ohosohate (NADP). All standards were purchased from Sionamide adenine dinucleotide ohosohate (NADP). All standards were purchased from Sionamide adenine dinucleotide ohosohate (NADP). All standards were purchased from Sionamide adenine dinucleotide ohosohate (NADP). All standards were purchased from Sionamide adenine dinucleotide of the standards were purchased from Sionamide adenine dinucleotide of the standards were purchased from Sionamide adenine dinucleotide (NAD).

ma. Deisenhofen, Germany.

5 TCA solution: 500 g trichloro-acetic acid, dissolved in H2O ad 2 I (25% w/v).

10 mM Tris buffer: 121.1 mg trishydroxymethyl aminomethane (Sigma, Deisenhofen, Germany) dissolved

in 100 ml H2O, titrated to pH = 10.4 with NaOH.

400 mg sulforhodamine B (Sigma, Deisenhofen, Germany), dissolved in 100 ml of 1%

(v/v) acetic acid.

K22-compounds were synthesised by the department of chemistry at Klinge Pharma

GmbH, Munich, Germany. Stock solutions: a 10 mM solution was prepared in dimethylsulloxide (DMSO) and stored at -18°C; further dilution steps were done in ethanol.

Cell line: The human-derived tumour cell line HeoG2 (liver carcinoma) was obtained from the

American Type Culture Collection (ATCC), Rockville, Maryland, USA.

Growth medium:

SRB solution:

Test substances:

[0040] Richter's Improved Minimal Essential Medium, Zino Option (INEM-ZO), was purchased from Gibco BRL. Life Technologie (Egganstein, Germany) (Richter, A., Santior, K.K. and Evans, V.I. (1972). J. Natl. Cancer Inst. 42 o 1705-1712). The medium powder was dissolved in delonised water, the pH Ittrated to 7.2 with Hcl I/NacOH and stell-rised by Ilfitation. The medium was supplemented with 8' och 10% feltad areum (FCS), PM Systems GmbH. Aldehbach, Germany; 100 µg/l Insulin (Boehringer, Mannheim, Germany) and 50,000 IU/I penicillin + 50 mg/l streptomycin (Sigma, Deisenbriden, Germany).

[0041] HEPES-butfored IMEM-ZO: This medium was used for incubation of HopG2 cells with the radiolabeled precursor. In contrast to the above-described Richter's IMEM-ZO, it did not contain niscinamice, NaHCO₃ and FCS. This medium was specifically prepared by Gibco BRL, Life Technologies (Eggenstein, Germany). The medium was butfered with 20 mM HEPES (Sigma, Deisenhofen, Germany) and the pH was adjusted to 7.2. The medium was sterilized by filtration.

50 Determination of NAD(P) synthesis from [14C]nlacinamide:

Cell culture:

[O042] The cells were detached from 75 cm² flasks by removing the growth medium and adding 3 ml tryssin/EDTA solution to each flask. After about 5 minutes incubation at 37°C, when the cells were detached from the surface of the dishes, tryseinization was stopped by adding 3 ml Richter's IMEN-ZO medium containing 10 % FCS. The cells were suspended by repeated pipetting. For prediution, an aliquot of 20µl was added to 10 ml Casyton isolonic solution (No. 043-90037P, Schaffe System, Puetlingen, Cemany) using a System Auto Dilutor Type AD-260 (Tips, Medical Elec-

tronics Co. Ltd., Kobe, Japan). The cell number was determined by electronic cell volume measurements and counting with a CASY 1 Cell Counter + Analyzer System, Model TTC (Schärfe System, Reutlingen, Germany) equipped with a 60 µm capillary. Following dilution in IMEM-ZO containing 10 % FCS, the cells were finally seeded at a density of 4x108/ 10 ml per sample in Ø 10 cm tissue culture dishes (Greiner, Frickenhausen, Germany) and incubated at 37°C in a humidified atmosphere of 5 % CO₂ in air.

[0043] After one day, when the cells were adherent to the dishes, the cultures were replenished with IMEM-ZO containing 5 % FCS plus the test compound or the vehicle. Concentrations of organic solvents in the medium after addition of the test substance did not exceed 0.1 % in any case. The cells were then incubated at 37°C for 17 hours in a humidified atmosphere of 5 % CO2 in air. This preincubation period is not necessary to achieve a distinct inhibitory action of the compounds and can be shortened for example to 2 or 0 hours. After this period of time, the medium was again discarded and 4 ml HEPES-buffered IMEM-ZO containing the test compound or the vehicle and 0.5 µCi/ml [14C] niacinamide were added to each culture for an additional 5 hours at 37°C and 100 % humidity. Just before the cells were harvested with a cell scraper and transferred to 15 ml polypropylene tubes, a 100 µl aliquot was taken from the incubation medium to determine the radioactivity. The culture dishes were rinsed with 4 ml saline supplemented with 10 mM nlacinamide and the solutions were pooled with the respective cell suspension. The cells were collected by centrifugation at 250 g for 5 minutes at 4°C.

Extraction of pyridine nucleotides

[0044] Pyridine nucleotides were extracted by a modification of the procedure of Chatteriee et al. (Chatteriee, S., Hirschler, N.V., Petzold, S.J., Berger, S.J. and Berger, N.A. (1989) Mutant Cells Defective in Poly(ADP-ribose) Synthesis due to Stable Alterations in Enzyme Activity or Substrate Availability. Exp. Cell Res. 184: 1-15). Briefly, each cell pellet was suspended in 200 µt ice-cold 0.5 M perchloric acid and incubated on ice for 20 minutes. After this period of time, the acid extracts were neutralized by adding 55 µl of a KCVKOH solution and centrifuged at 2500 g for 10 minutes at 4°C. Supernatants were collected and stored at 20°C until separation by chromatography. A 10 ut aliquot was taken from each supernatant to measure the total amount of radioactivity in the cell extract.

Thin-layer chromatography

[0045] The 14C-labeled components of the cell extracts were separated and identified using two thin-layer chromatography (TLC) systems. 2 µl of each cell extract was transferred to a cellulose and a poly(ethyleneimine) (PEI) cellulose TLC foil using a DC-Probenautomat III (CAMAG, Muttenz, Switzerland). The cellulose foils were developed using 1 M NH4 acetate:ethanol (3:7) as solvent (Pinder, S., Clark, J.B. and Greenbaum, A.L. (1971) The Assay of Intermediates and Enzymes Involved in the Synthesis of the Nicotinamide Nucleotides in Mammalian Tissues, Methods in Enzymology. Academic Press, New York. Vol. XVIIIB pp. 20-46). The PEI cellulose plates were developed with 0.05 M lithium chloride (Barton, R.A., Schulman, A., Jacobson, E.L. and Jacobson, M.K. (1977) Chromatographic Separation of Pvridine and Adenine Nucleotides on Thin Layers of Poly(ethyleneimine) Cellulose. J. Chromatogr. 130: 145-150). [0046] The chromatograms were run with added non-radioactive standards of NAD, NADP, NAM, dNAM, dNAD. niacin and niacinamide, and the spots were identified by UV absorption. See Table 4 for R_F values. Results are expressed as mean ± S.D. For autoradiography, the chromatograms were exposed to an imaging plate BAS-IIIs (Fuji

Photo Film Co., Ltd., Japan) in a hypercassette (Amersham Buchler GmbH & Co. KG, Braunschweig, Germany) for at least two days. To avoid high background activity, the cassette was placed in a lead box. After exposure, the imaging plate was read in the bio-imaging analyzer FUJIFILM BAS-1500 (Fuji Photo Film Co., Ltd., Japan), The portion of each [14C]-labeled component in the cell extracts was determined as percentage of total radioactivity with the software TINA

2.0 (raytest isotopenmessgerate GmbH, Straubenhardt, Germany).

Table 4

Standard	Matrix / Solvent		
	PEI cellulose / LiCI	Cellulose / NH ₄ acetate:ethanol	
Niacin	0.45 ± 0.03 (n = 12)	0.73 ± 0.04 (n = 14)	
Niacinamide	0.77 ± 0.01 (n = 14)	0.80 ± 0.04 (n = 14)	
dNAM	0.18 ± 0.04 (n = 4)	0.19 ± 0.04 (n = 4)	
NAM	0.52 ± 0.08 (n = 3)	0.19 ± 0.05 (n = 3)	
dNAD	0.07 ± 0.01 (n = 10)	0.09 ± 0.01 (n = 10)	

Table 4 (continued)

Standard	Matrix / Solvent			
	PEI cellulose / LiCl	Cellulose / NH ₄ acetate:ethanol		
NAD	0.37 ± 0.02 (n = 14)	0.11 ± 0.02 (n = 14)		
NADP	0.02 ± 0.01 (n = 13)	0.04 ± 0.01 (n = 13)		

[0047] The amount of ¹⁴C-labeled derivatives was calculated by multiplying the total radioactivity of the cell extract by the percentage recovered in each derivative. The results of the assays as shown in Table 3 above are expressed as pmot [14C]ND(P) per 10⁸ cells and as pmot [14C]ND(P) per mg protein. Cell count and cellular protein were determined from cultures prepared in parallel without radioactive precursors.

Protein determination:

[0048] The cellular protein was determined with the bicinchoninic acid (BCA) assay purchased from Pierce, Rockford, IL, USA, according to the manufacturer's instructions. The colour of the samples produced from the reaction was measured spectrophotometricating (COBAS FARM IF. F. Hoffmann-La Roche AG, Basel, Switzfand).

Determination of cell growth under high-density conditions: Cell culture

[0049] The cells were detached from 75 cm2 flasks by removing the growth medium and adding 3 ml trypsin/EDTA collution to seak will. After 5 mlutules inclustion at 37°C, when the cells were detached from the surface of the dishes, trypsinisation was stopped by adding 3 ml Richter's IMEM-ZO medium containing 10 % FCS. The cells were suspended by repeated piperting. For prediction are aligned of 20 µl was added to 10 ml cells approximations coloring (No. 043-90037P, Schaffe System, Reutingen, Germany) using a Systems Auto Dilutor Typs AD-280 (Toe, Medical Electronics Co. Ltd., Kobe, Japan). The cell number was determined by electronic cell volume measurements and counting with a CAST (Cell Counter 4-radyser System, Model TTC (Eschaffe System, Reutingen, Germany) equipped with a 60 µm capillary.

Following dilution in growth medium the cells were finally seeded at a density of 200,000 cells per ml and well in 24-well culture dishes (Griene, Frickenhausen, Germany). Additionally, three negative control wells were incubated in growth medium without any cells.

[0050] After one day, when he cells were achieven to the delhes, the cultures were replenished with fresh medium containing 5.% reconstructions of the velocity of the velocit

Sulforhodamine B assay (SRB)

[0051] Determination of cell growth was performed by unspecific protein staining with sulforhodamine B according to Skehan et al. (Skehan, P. et al. (1990) New Colorimetric Cytotoxicity Assay for Anticancer-Drug Screening. J. Natl. Cancer inst. 82: 1107-1112).

[0052] The drug incubation period of the cells was stopped by the addition of 250 µl of Lee cold TCA solution into the growth medium. After one hour incubation in the refrigerant, the superinated was discarded, the dishes were resed five times with delonised water, dried at room temperature (RT) and finally stored in the refrigerator until staining. 0.5 ml of SRB solution was appleted into each well and incubated at room temperature for 30 minutes, thereafter, the staining solution was decarted, the dishes were washed four times with 1% (v/l) acetic acid and dried again at RT. SRB stain unspecifically bound to protein was released by adding 1 ml of 10 mM Tris buffer per well and gentle shaking for 5 minutes. 100 µl aliquoted to each well were transferred to a 95-wind incrotiter plate and the light sottriction at 515 nm wavelength was read in an ELISA-reader (Bio-Teix, Dedux, Gödenschof, Germany). The mean value of negative control wells was subtracted from the test asmiller readings.

THERAPEUTIC ADMINISTRATION FORMS

[0053] The production of medicaments with an amount of one or more compounds according to the invention and/ or their use in the application scording to the invention occurs in the usubnary amenance by means of common pharmaceutical technology methods. For this, the active ingredients as such or in the form of their salts are processed together with suitable, pharmaceutically acceptable adjuvents and carriers to medicinal forms suitable for the various

indications and types of application. Thereby, the medicaments can be produced in such a manner that the respective desired release rate is obtained, for example a quick flooding and/or a sustained or depol effect.

[0054] Preparations for parenteral use, to which injections and infusions belong, are among the most important systemically employed medicaments for tumor treatment as well as for other indications.

- 5 [0055] Preferably, injections are administered for the treatment of tumors. These are prepared either in the form of vials or also as so-called ready-to-use injection preparations, for example as ready-to-use syringes or single use syringes in addition to perforation bottles for multiple withdraweak. Administration of the injection preparations can occur in the form of subcutaneous (s.c.), intramuscular (i.m.), intravenous (i.v.) or intracutaneous (i.c.) application. The respective suitable injection forms can especially be produced as solutions, crystal suspensions, nanoparticular or colloid-dispress systems, such as for example, hydrosols.
 - [0056] The injectable formulations can also be produced as concentrates which can be adjusted with aqueous isotonic dilution agents to the desired active ingredient dosage. Furthermore, they can also be produced as powders, such as for example lyophilisates, which are then preferably dissolved or dispersed immediately before application with suitable dilutents. The infusions can also be formulated in the form of isotonic solutions, tat emulsions, liposome formulations, microemulsions and liquids based on mixed micells, for example, besed on phospholylids. As with injection proparations, infusion formulations can also be prepared in the form of concentrates to dilute. The injectable formulations can also be applied in the form of continuous infusions as in stationary as well as in outpatient therapy for example in the form of mini-toward.
- [0057] Albumin, plasma expanders, surface active compounds, organic solvents, pH influencing compounds, compared plex forming compounds or polymeric compounds can be added to the parenterial medicinal forms, especially as substances for influencing the adsorption of the active ingredients to protein or polymers or also with the aim of decreasing the adsorption of the active ingredient to materials such as injection instruments or packaging materials, for example plastic or class.
- [0058] The active ingredients can be bound to nanoparticles in the preparations for parenteral use, for example on finely dispersed particles based on polymetrih(archieus) polyacetates, polygipodates, polyamino acids or polyather urethanes. The parenteral formulations can also be constructively modified as depot preparations, for example on the multiple unit principle, where the active ingredients are incorporated in a most finely distributed and/or dispersed, suspended form or as crystal suspensions, or on the single unit principle, where the active ingredient is enclosed in a medicinal form, for example, a tablet or a seed which is subsequently implanted. Chort, these implantations or depot andigeneers to include the calculations.
- medicinal form, for example, a tablet or a seed which is subsequently implanted. Often, these implantations or depot medicaments in single unit and multiple unit medicinal forms consist of so-called floodegradable polymers, such as for example, polyether uerthenes of lactic and glycotic acid, polyether urethenes, polyamino acids, polyfinethjacry/ates or polysaccharides.
- [0059] Sterilized water, pH value influencing substances, such as for example organic and inorganic acids or bases as well as their salts, buffer substances for setting the pH value, agents for isotonicity, such as for example sodium of chief, monosodium, carbonate, glucose and fructose, tensides and/or surface active substances and emulatiliers, such as for example, partial fatty acid esters of polyoxethylene (Cremophore), fatty olds such as for example permut oil, solybean oil and castor oil, synthetic fatty acid esters, such as for example permut oil, solybean oil and castor oil, synthetic fatty acid esters, such as for example permut oil, solybean oil (Mighyot8) as well as polymer adjuvents such as for example getter, distrant, polyviny/prointion, organic solvent additives which increase solutility, such as for example proylene glycol, ethanol, NN-dimethylacetamide, propylene glycol or complex forming compounds such as for example citates and urea, preservatives, such as for example sodium sutilitie and stabilizers, such as for example EDTA, are suitable as additivents and carriers in the production of preparations for parenteral use.
- [0060] In suspensions, addition of thickening agents to prevent the settling of the active ingradients from tensides and poptizes, to secure the ability of the sediment to be shaken, or complex formers, such as EDTA, ensuess. This can also be achieved with the various polymeric agent complexes, for example with polyethylene glycols, polystyrol, carboxymethyleulluses, Plurcaries® or polyethylene glycol sorbitan fatly acid esters. The active ingredient can also be incorporated in liquid formulations in the form of inclusion compounds, for example with cyclodaxtins. As further acid provides, dispension agents are also suitable. For production of lyophilisates, builders are also used, such as for example or mannile, detarns, asocharose, human albumin, lactose, PVP or polatin variation.
 - [0061] As long as the active ingredients are not incorporated in the liquid medicinal formulations in the form of a base, they are used in the form of their acid addition salist, Pytrates or solvates in the preparations for parenteral use. [0062] A further systemic application form of importance is percral administration as tablets, hard or soft gelatin capsules, coaled tablets, powders, pollets, microcapsules, obling compressives, granules, chewable tablets, lozenges, gume or socknist. These sold percral administration forms can also be prepared as sustained action and/or depot systems. Among these are medicaments with an amount of one or more micronized active ingredients, diffusions and erosion forms based or matrices, for example by using fats, war-like and/or polyment compounds, or so-called res-rovior systems. As a relating agent nativor agent to controlled release, film or matric froming substances, such as

for example ethylocellulose, hydroxypropylmethylocellulose, polymeth)acrylate derivalives (for example Eudragit®), nydroxypropylmethylocellulose phthalate are suitable in organic solutions as well as in the form of aqueous dispersions. In this connection, so-called bio adhesive preparations are also to be named in which the increased retention time in the body is achieved by intensive contact with the mucus membranes of the body. An example of a bio-adhesive solvmer is the aroup of Carbonness.

[0053] For sublingual application, compressives, such as for example non-disintegrating tablets in obtory from of a suitable size with a solve release of active ingredient, are especially suitable. For purposes of a targeted release of active ingredients in the various sections of the gastrointestinal tract, mixtures of pellets which release at the various places are employable, for example mixtures of gastric fluid soluble and small intestine soluble and/or gastric fluid resistant and large intestine soluble pellets. The same goal of releasing at various sections of the gastrointestinal tract can also be conceived by suitably produced laminated tablets with a core, whereby the coating of the agent is quickly released in gastric fluid and the core of the agent is solwy released in the small intestinal milliou. The goal of controlled release at various sections of the gastrointestinal tract can also be attained by multitayer tablets. The pellet mixtures with differentiaty released ingred can be filled into hard gelatin casquality released and reach as the filled into the deplate in casquality released and can be filled into the deplate in casquality.

[0054] Anti-stick and lubricant and separating agents, dispersion agents such as llame dispersed silicone dioxide, disintegrants, such as various starch types, PVC, cellulose esters as granulating or retaining agents, such as for example wax-like and/or polymeric compounds on the basis of Eudragit®, cellulose or Cremophor® are used as a further adjuvents for the production of compressives, such as for example tablets or hard and soft gelatin capsules as well as coated tablets and oranulates.

[0065] Anti-cxxtants, sweetening agents, such as for example saccharose, sylle or mannile, masking flavors, arcmatics, presenvieves, colorants, buffer substances, direct tabeling agents, such as for example microcrystalline celulose, starch and starch hydrolysates (for example Colutab9), actose, polyethylene glycols, polyvinylymotidoria and dicactium phosphate, lubricants, filliors, such as lactose or starch, binding agents in the form oil factose, starch varieties, such as for example wheat or core and/or rice starch, cellubose derivatives, for example methylecilliose, hydroxypropy/collulose or silice, lateum powder, stearates, such as for example magnesium stearate, aluminum stearate, calcium stearate, taic, elicionized tale; clearte acid, seely tachor) and hydrated falts are used.

[0066] In this connection, or altherapeutic systems constructed especially on osmotic principles, such as for example GIT (pastrointestinal therapeutic system) or OROS (or all osmotic system), are also to be mentioned.

[0067] Ellevaseont labels or tabs. both of which represent immediately drinkable instant medicinal forms which are or quickly disavoled or suspended in water are among the percently administratible compressives. Among the percently administratible compressives. Among the percently administratible forms are also solutions, for example drops, juices and suspensions, which can be produced according to the two types given method, and can still contain preservatives for increasing stability and optionally amonatics for reasons of easier intake, and colorants for better differentiation as well as antioxidants and/or vitamins and sweeteners such as sugar or artificial sweetening agents. This is also true for inspisated juices which are formulated with water 35 before ingestion, ion exchange resins in combination with one or more active ingredients are also to be mentioned for the production of fould incosebable forms.

[0068] A special release form consists in the preparation of so-called floating medicinal forms, for example based on tablets or pellets which develop gas after contact with body fluids and therefore felos on the surface of the gastric fluid. Furthermore, so-called electronically controlled release systems can also be formulated by which active ingredient releases can be selectively adjusted to individual nearly.

[0069] A further group of systemic administration and also optionally lopically effective medicinal forms are represented by rectally applicable medicaments. Among these are suppositories and enema formulations. The enema formulations can be prepared based on tablets with aqueous solvents for producing this administration form. Rectal capsules can also be made available based on gelatin or other carriers.

[0070] Hardened fat, such as for example Witepsol®, Massa Estarinum®, Novata®, coconut fat, glycerol-gelatin masses, glycerol-soap-gels and polyethylene glycols are suitable as suppository bases.

[0071] For long-term application with a systematic active ingredient release up to several weeks, pressed implants are suitable which are preferably formulated on the basis of so-called biodegradable polymers.

[0072] As a further important group of systemically active medicaments, transdemal systems are also to be emphasized which distinguish themselves, as with the above-mentioned rectal forms, by circumventing the liver circulation system and/or liver metabolism. These plasters can be especially prepared as transdemal systems which are capable of releasing the active ingredient in a controlled menner over longer or shorter time periods based on different layers and/or mixtures of suitable adjuvents and carriers. Aside from suitable adjuvents and carriers such as solvents and polymeric components, for example based on Eudragit®, membrane infiltration increasing substances and/or permesition promoters, such as for example older acid, Azone®, displine acid derivatives, shand, urea, propriy[opcid are suitable in the production of transdemal systems of this type for the purpose of improved and/or accelerated penetration.

[0073] As topically, locally or regionally administration medicaments, the following are suitable as special formula-

tions: vaginatily or genitally applicable emulsions, creams, foam tablets, depot implants, ovular or transurethral adminstration installation solutions. For opthalmological application, highly sterile eye ointments, solutions and/or drops or creams and emulsions are suitable.

- [0074] In the same manner, corresponding oblogical drops, ointhents or creams can be designated for application to the ear. For both of the above-mentioned applications, the administration of semi-coid formulations, such as for sxample gels based on Carbopole® or other polymer compounds such as for example polyviny(pyrolidone and cellulose derivalives) is also possible.
- [0075] For customary application to the skin or also to the mucus membrane, normal emulsions, gels, ointments, creams or mixed phase and/or ampliphilic emulsion systems (olivater-veatroir) imbed phase) as well as lipsonnes and transfersomes can be named. Sodium algenate as a gel builder for production of a suitable foundation or celluloses derivatives, such as for example gua or xamibene gum, inorganic gel builders, such as for example aluminum hydroxides or bentonities (so-called thiootopic gel builder), polyacrylic acid derivatives, such as for example Carbopol®, polyarrylpyrolidone, microcrystalline cellulose or carboxymethylcellulose are suitable as adjuvents and/or carriers. Furhermore, amphiphilic low and hish molecular weight compounds as well as phospholigida era suitable. The gels can be present either as hydrogels based on water or as hydrophobic organogels, for example based on mixtures of low and hish molecular varafilir hivdocarbons and vassiline.
- [0076] Anionic, cationic or neutral tensides can be employed as emulatifiers, for example alkalized scaps, methyl sospe, sufferance for compounds, cationic scaps, high fally alcholes, partial falty aid elsters, of softial rand polyoxyeltylene sorbitan, for example lanette types, wool wax, lanolin, or other synthetic products for the production of bilwater faults with softial results and polyoxyeltylene sorbitan, for example lanette types, wool wax, lanolin, or other synthetic products for the production of bilwater faults with softial results and softial results and the production of bilwater faults.
- [0077] Hydrophilic organogals can be formulated, for example, on the basis of high molecular polyethylene glycols. These gel-like forms are weak-blow. Vasalien, actual or symbritic waxes, talty eachs, latty alchols, fatty acide starts, for example as monor, dr. or triglycerides, paraffin oil or vegetable oils, hardened castor oil or occount oil, pig fatt, symbretic fats, or example based on acrylic, apprintle, subtra and stearine acids, such as for example Sottamore or triglyceride mixtures such as Miglyol® are employed as lipids in the form of fat and/or oil and/or wax-like components for the production of oil minnests, creams or emulsions.
- [0078] Osmotically effective acids and bases, such as for example hydrochloric acid, citric acid, sodium hydroxide solution, potassium hydroxide solution, monosodium carbonate, further buffer systems, such as for example citrate, phosphate. This buffer or triethanolamine are used for adjustion the DH value.
- [0079] Preservatives, for example such as methyl- or propyl benzoate (parabenes) or sorbic acid can be added for increasing stability.
 - [0080] Pastes, powders or solutions are to be mentioned as further topically applicable forms. Pastes often contain lipophilic and hydrophilic auxiliary agents with very high amounts of fatty matter as a consistency-giving base.
 - [0081] Powders or topically applicable powders can contain for example starch varieties such as wheat or rice starch, flame dispersed silicon dioxide or silica, which also serve as diluents, for increasing flowability as well as fubricity as well as for preventing accidence rates.
 - [0082] Nose drops or nose sprays serve as nasal application forms. In this connection, nebulizers or nose creams or ointments can come to use.
- [0083] Furthermore, nose spray or dry powder formulations as well as controlled dosage aerosols are also suitable
 40 for systemic administeration of the active ingredients.
- [0049] These pressure and/or controlled dosage aerosols and dry powder formulations can be inhaled and/or insuflated. Administration forms of this type also certainly have importance for direct, regional application in the lung or bronchi and larynx. Thereby, the dry powder compositions can be formulated for example as active ingredient-soft pellets, as an active ingredient-pollet initiature with suitable acriners, such as for example lactose and/or glucose. For finishation or insufficiation, common applications are suitable which are suitable for the treatment of the nose, mouth and/or pharynx. The active ingredients can also be applied by means of an ultrasonic nebulizing device. As a propellant gas for aerosol spray formulations and/or controlled dosage aerosols, laterillucorothane or FIFC 124 and of heptaflucorpropane or HFC 227 are suitable, wherein non-flucinisted hydrocarbons or other propellants which are gaseous at normal pressure and core interpretative, such as for example process. Butter or dimethy ether can be preferred.
- o Instead of controlled dosage aerosols, propellant-free, manual pump systems can also be used. [0085] The progellant gas aerosols can also suitably contain surface active adjuvents, such as for example isopropyl myristate, colvoxyethylene sorbitan fatty acid ester, sorbitan fatty acid ester.
 - [0086] For regional application in situ, solutions for installation, for example for transurethral administration in bladder tumors or central tumors, or for profusion in liver tumors or other organizations are suitable.

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[0087]

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Claims

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- Biologically active substance which inhibits the cellular formation of niacinamide mononucleotide.
 - Biologically active substance according to claim 1 having an inhibitory activity on cellular NAD biosynthesis from the precursor niacinamide at concentrations of ≤ 10 µM of 50 %, preferably 80 %, most preferably 90 %.
- 3. Biologically active substance according to claim 1 or 2 which is an inhibitor of niacinamide phosphoribosyl transferase.
 - Biologically active substance according to claim 1 or 2 which inhibits the transport of niacinamide across the cell membrane.
 - 5. Biologically active substance according to claim 1 or 2 which is selected from:
 - 1-[4-(1-benzhydryl-piperidine-4-yl)-butyl]-3-pyridine-3-yl-urea;
 - 4-benzhvdryl-piperazine-1-carboxylic acid-[6-(3-pyridine-3-yl-methylureido)-hexyl]-amide;
 - 1-(3,3-diphenylpropyl)-3- [6-(3-pyridine-3-yl-methylureido)-hexyl]-urea;
 - 1-[5-(1-benzhydryl-piperidine-4-yl)-pentyl]-3-pyridine-3-yl-thiourea;
 - 6-(4-benzhydryl-piperazine-1-yl)-hexanoic acid-(2-pyridine-3-yl-ethyl)-amide;
 - 1-(6,6-diphenyl-5-hexenyl)-3-(pyridine-3-yl-methylene-amino)- thiourea:

- N-(4-{1-{4-(1-benzhydryl-piperidine-4-yi)-butyl}-piperidine-4-yi]-butyl)-3-pyridine-3-yl-propanoic acid amide; 1-{4-(1-benzhydryl-piperidine-4-yi)-butyl}-3- (2-pyridine-3-yl-ethyl)-urea;
- N-[2-[5-(4-benzhydryl-piperazine-1-yl-methyl)-1-methyl-1H-pyrrole-2-yl]-ethyl]-3-pyridine-3-yl-acrylamide:
- N-[2-[5-[4-0enZhydryl-piperazine-1-yl-methyl]-1-methyl-1H-pyrrole-2-yl]-ethyl]-3-pyridine-3-yl-acrylamide 1-[4-[1-(naphthalin-2-sulfonyl)-piperidine-4-yl]-butyl]-3-pyridine-3-yl-urea;
- 1-[4-[1-(10,11-dihydro-dibenzene[b,f]azepine-5-carbonyl)-piperidine-4-yl]-butyl]-3-pyridine-3-yl-urea; and 2-amino-3-(4-hydroxy-3-(2-(4-[4-(3-pyridine-3-yl-acryloyl-amino)-butyl]-piperidine-1-carbonyl]-pnenylazo)-phenyl
- Pharmaceutical composition comprising a biologically active substance according to any of the claims 1 to 5, or a pharmaceutically acceptable salt thereof, optionally together with pharmaceutically acceptable formulation additive.
 - 7. Pharmaceutical composition according to claim 6 for the treatment of cancer or immunosuppression in mammals.
- Pharmaceutical composition according to claim 7, wherein the cancer is selected from breast, prostate, lung, colon, cervix, ovary, skin, CNS, bladder, pancreas and leukemia and lymphoma.
 - Pharmaceutical composition according to any of the claims 6 to 8 which is formulated for intraperitoneal, subcutaneous, oral, intravenous, rectal, buccal, intramuscular, intravaginal, topic or pulmonal administration.
 - 10. Use of a biologically active substance according to any of the claims 1 to 5, or a pharmaceutically acceptable salt thereof, for the preparation of a pharmaceutical composition for the treatment of cancer in mammals.
- Method for screening and detecting biologically active substances according claims 1 to 5 comprising:

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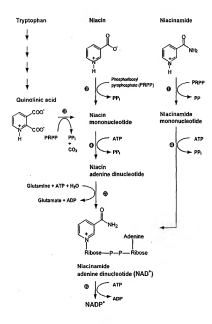
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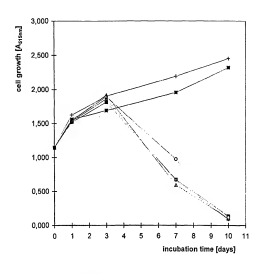
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- incubating cultured cells selected from HepG2 cells, U.87 MG cells, MCF7 MI cells, Caki-1 cells, HL-60 cells, PC3 cells, U-97 MG cells, ASG cells and KG-1 cells in the presence of 1⁴Cylactamids and a substance to be tested for its activity to inhibit the cellular formation of niacinamide mononucleotide, effecting lavia of the cells:
- isolating and separating the [14C]-labelled compounds and measuring the amount of formed labelled niacinamide mononucleotide. NAD and NADP.
 - 12. Method for determining the dependency of a cell type on niacinamide as a precursor for NAD synthesis comprising: incubating cells to be assayed in the presence of a substance according to claims 1 to 5 in a medium containing only niacinamide as a NAD synthesis precursor, and performing a cytotoxicity assay after the incubation period.



- niacinamide phosphoribosyl transferase
- 9 niacin phosphoribosyl transferase
- 9 quinolinic acid pyrophosphate phosphoribosyl transferase
- NAD pyrophosphorylase
- NAD synthetase
- O NAD kinase

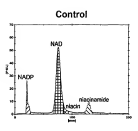
Figure 1: Biochemical Pathways of NAD(P)* Biosynthesis



Test Articles:

-■- K22.339 1x10-5 M
-△- K22.339 3x10-6 M
-○- K22.339 3x10-7 M
-- K22.339 3x10-7 M
-- K22.339 3x10-8 M
-=- Control

Figure 2: Induction of "delayed cell death" in HepG2 cells.



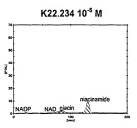


Figure 3: Inhibitory action of K22.234 on NAD(P) biosynthesis from [\$^1C]niacinamide in HepG2 cells. The radioactive metabolites of the cell extracts were separated on PEI cellulose using 0.05 M LICI as solvent.



European Patent Office

PARTIAL EUROPEAN SEARCH REPORT

which under Rule 45 of the European Patent Convention EP 99 10 3814 shall be considered, for the purposes of subsequent proceedings, as the European search report

DOCUMENTS CONSIDERED TO BE RELEVANT Category : Gitation of document with indication, where appropriate, of relevant passages Relevant to clarm CLASSIFICATION OF THE APPLICATION (INLCL?) WD 97 48696 A (REITER FRIEDEMANN ;HASMANN MAX (DE); LOESER ROLAND (DE); BIEDERMAN)
24 December 1997 (1997-12-24)
4 Table 1; claims 18-40 * C07D213/24 C07D213/38 C070213/53 C07D213/89 C07D401/12 WO 97 48397 A (KLINGE CO CHEM PHARM FAB ;BIEDERMANN ELFI (DE); HASMANN HAX (DE);) 24 December 1997 (1997-12-24) A61K31/44 5-10 A61K31/495 A61K31/55 * Tables 1 and 2; claims 1-18 * C1201/68 TECHNICAL FIELDS CO7D A61K C120 INCOMPLETE SEARCH The Search Division comilders that the present application, or one or more of its claims, does/do not comply with the EPC to such an extent that a manningle search into the state of the art cannot be searched unty critically, for these claims. Claims searched completely: Claims searched incompletely: Clares and assembled Resson for the limitation of the search: see sheet C Place of search Date of compression of the search MUNICH 6 September 1999 Uiber, P CATEGORY OF CITED DOCUMENTS T: theory or principle underlying the invention E: earlier patient document, but published on, or after the fling date D: document offed in the explication L: document offed for other reasons & ; member of the same petent family, corresponding



INCOMPLETE SEARCH SHEET C Application Number EP 99 10 3814

Claim(s) searched completely: 5.11.12

Claim(s) searched incompletely: 1-4,6-10

Reason for the limitation of the search:

Present claims 1-4 and relate to a compound or composition or defined by reference to a desirable characteristic or property, namely:

- inhibition of the cellular formation of miacinamide mononucleotide

The claims cover all compounds or compositions or uses having this characteristic or property, whereas the application provides support within the meaning of Article 84 EPC and/or disclosure within the meaning of Article 83 EPC for only a very limited number of such compounds or compositions or uses. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claims sociated a territy thricine as EPC). An interpt is read to the claim sociated a territy thricine as EPC). An interpt is a supported to the product compound by reference to a result to be achieved, Again, this lack of clarity in the present case is such as to render a meaningful search over the whole of the claimed scope impossible. Consequently, the search has been carried out for those parts of the claims which appear to be clear, supported and disclosed, namely those parts relating to the compounds of claim 5. compositions containing the compounds of claim 5. on seas of the compounds of claim 5 according to claim 10 and the methods of claim 1 and 12.

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 99 10 3814

This arries talls the palent family members relating to the patent obsciments oded in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Educ

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b For more details about this annex ; see Official Journal of the European Patent Office, No. 12/82